Opsin-based photopigments expressed in the retina of a South American pit viper, Bothrops atrox (Viperidae)

Christiana Katti,1,2 Micaela Stacey-Solis,1 Nicole A. Coronel-Rojas,1 and Wayne I.L. Davies3–6

1Escuela de Ciencias Biológicas, Pontificia Universidad Católica del Ecuador, Apartado 17-01-2184, Quito, Ecuador; 2Programa Becas Prometeo, Secretaría Nacional de Educación Superior, Ciencia, Tecnología e Innovación de la República del, Quito, Ecuador; 3Oceans Graduate School, University of Western Australia, Perth, WA 6009, Australia; 4Oceans Institute, University of Western Australia, Perth, WA 6009, Australia; 5School of Biological Sciences, University of Western Australia, Perth, WA 6009, Australia and 6Lions Eye Institute, University of Western Australia, Perth, WA 6009, Australia

Abstract

Although much is known about the visual system of vertebrates in general, studies regarding vision in reptiles, and snakes in particular, are scarce. Reptiles display diverse ocular structures, including different types of retinae such as pure cone, mostly rod, or duplex retinas (containing both rods and cones); however, the same five opsin-based photopigments are found in many of these animals. It is thought that ancestral snakes were nocturnal and/or fossorial, and, as such, they have lost two pigments, but retained three visual opsin classes. These are the RH1 gene (rod opsin or rhodopsin-like-1) expressed in rods and two cone opsins, namely LWS (long-wavelength-sensitive) and SWS1 (short-wavelength-sensitive-1) genes. Until recently, the study of snake photopigments has been largely ignored. However, its importance has become clear within the past few years as studies reconsider Walls’ transmutation theory, which was first proposed in the 1930s. In this study, the visual pigments of Bothrops atrox (the common lancehead), a South American pit viper, were examined. Specifically, full-length RH1 and LWS opsin gene sequences were cloned, as well as most of the SWS1 opsin gene. These sequences were subsequently used for phylogenetic analysis and to predict the wavelength of maximum absorbance ($A_{max}$) for each photopigment. This is the first report to support the potential for rudimentary color vision in a South American viper, specifically a species that is regarded as being nocturnal.

Introduction

The visual system of vertebrates has evolved over millions of years to adapt to diverse habitats with varying spectral ranges and intensities of light. As such, many animals have developed activities that are generally restricted to particular times of the day (e.g., nocturnality, crepuscularly, or diurnality) or in response to a specific ecological niche (e.g., burrowing/fossorial or arboreal behaviors) (reviewed in Davies et al., 2012 and Gerkema et al., 2013). Reptiles, like many vertebrates, possess different types of retinae. Based on photoreceptor morphology, most reptilian retinas appear to be duplex (consisting of both rods and cones), but they can vary from being rod dominated to retinae that consist of pure cones (Janke & Arnason, 1997; Loew & Govardovskii, 2001). Generally, photoreceptor morphological diversity appears to be due to adaptive evolution of the eye to different light environments within a diverse array of habitats; however, the visual systems of reptiles (and related sister groups) are further complicated by the potential for “transmutation.”

Postulated by Walls throughout the 1930s and 1940s, the “transmutation theory” was developed to explain the rod-like morphology of cones (and vice versa) present in the retinas of many reptiles (Walls, 1934, 1942). Specifically, Walls argued that at some point during the evolution of early tetrapods, many ancestral forms shifted their lifestyles from nocturnality to diurnality, most likely to take advantage of the sun for warmth and increased cellular metabolism, especially since these poikilotherms were unable to regulate intrinsic body temperatures (Gerkema et al., 2013; Walls, 1942). As such, they depended solely on photopic vision, which resulted in the partial or complete loss of rods. With the evolution of predators, such as birds and mammals, early reptilian tetrapods were forced to become nocturnal to increase their chance of survival. To be able to detect low levels of light, these species had to modify their remaining photoreceptors (i.e., cones) to functionally act as rods. As a result, morphological changes occurred such that these photoreceptors transmuted into cells that...
either still resembled cones (but were sensitive to low intensities of light), or in most cases became a cone–rod hybrid to maximize photon capture (Walls, 1934, 1942).

Snakes are generally divided into two major infraorders: Scolecophidia (blind snakes and thread snakes) and Alethinophidia (all other snakes). Alethinophidia can be further separated into two superfamilies: Henophidia, which includes boids (pythons and boas), and Caenophidia, which is far more diverse and includes vipers, elapids, and colubrids (Hsiang et al., 2015; Scanlon & Lee, 2011). In general, scolecophidians are fossorial with reduced eyes that are covered with scales in some cases. Their visual systems have not been studied extensively; nonetheless, the few reports that are published suggest that scolecophidians have pure rod retinas (Simões et al., 2015). Henophidia includes many nonvenomous, crepuscular/nocturnal, and/or burrowing snakes (Collins & Conant, 1998) and is considered the more primitive superfamily of the infraorder Alethinophidia thus, it is believed that the visual system of boids and pythons is ancestral to that of the more evolutionarily advanced caenophidians (Crescitelli, 1972). Henophidians are primarily nocturnal, but can be active during the day, and therefore have duplex retinas that are rod dominant (90% rods vs. 10% cones). Caenophidia is a much more diverse group and includes various diurnal and nocturnal snakes; therefore, it is thought that the retinal composition of these species should be greatly variable in response to the their different visual requirements; for example, diurnal colubrids have pure-cone retinas compared to rod-based vision in some nocturnal species such as vipers (Hauzman et al., 2014).

Most reptiles possess five types of opsins-based photopigments (where the opsins protein is linked to a photosensitive chromophore) that are housed within each retinal photoreceptor: rod opsins (rhodopsin-like-1, RH1) is exclusively expressed in rods, with four different cone opsins classes (Bowmaker, 2008; Davies et al., 2012; Davies et al., 2009b; Yokoyama, 2000). The cone photopigments are composed of short-wavelength-sensitive-1 (SWS1), short-wavelength-sensitive-2 (SWS2), rhodopsin-like-2 (RH2), or long-wavelength-sensitive (LWS) opsins proteins. Unlike in other reptiles, the few studies conducted on snake vision suggest that scolecophidians only express the RH1 gene (Simões et al., 2015), whereas more advanced snakes possess SWS1, LWS, and RH1 opsins genes (Davies et al., 2009b;Emerling, 2017; Schott et al., 2016; Simões et al., 2016a; Simões et al., 2016b).

The aim of this study is to expand knowledge of the visual system of snakes by investigating the complement of visual pigments expressed in the retina of Bothrops atrox (the common lancehead) and predict their wavelength of maximum absorbance ($\lambda_{\text{max}}$). In particular, this species was chosen as B. atrox is a nocturnal, terrestrial viper (Viperidae) that is widely distributed in Central and South America, known to climb trees in search of prey, and is an adept swimmer (Stocker & Barlow, 1976). More specifically, the common lancehead inhabits a variety of habitats, including low montane humid forests, savannas, and rainforests. Indeed, this species is generally associated with the presence of water, with a preference for humid environments near streams, lakes, or wetlands (Campbell & Lamar, 2004; Oliveira & Martins, 2001), and plays an important ecological role in tropical rainforests. Notably, it is one of the most poisonous snakes of the region (Hussain et al., 2011). In general, scolecophidians only express the RH2 gene (Simões et al., 2015), whereas more advanced snakes possess SWS1, LWS, and RH1 opsins genes (Davies et al., 2009b;Emerling, 2017; Schott et al., 2016; Simões et al., 2016a; Simões et al., 2016b).

The results of this study showed that the common lancehead expresses three photopigment genes that were phylogenetically determined to be LWS, SWS1, and RH1. In addition, spectra were predicted from the complement of known tuning sites, which showed that this species is sensitive to wavelengths that range from the ultraviolet (UV) to the long-wavelength end of the visible spectrum, and that functionally resembles the more ancient “lower order” henophidian snakes than the “higher order” caenophidians, of which B. atrox is a member. Given that B. atrox is nocturnal and dwells in tropical rainforests, where light is extremely restricted, it was expected that only rods expressing RH1 would be identified. As such, the presence of cone photopigments is biologically intriguing. The study of the visual system of this nocturnal South American snake is important as it is the first viper from the region to be investigated, as well as being one of the first to research vision of any vertebrate in Ecuador.

Materials and methods

Animals and collection of samples

An adult South American Bothrops atrox (common lancehead) snake (Fig. 1) was collected from the Zamora-Chinchipe Province, Ecuador. Since B. atrox is a relatively difficult serpent to find and collect, due to various reasons such as its activity patterns, its aggressive behavior, as well its highly cryptic color pattern, only the eyes from one specimen, which was donated to perform this study, were used. This specimen was donated by and is registered in the Herpetology Museum of Pontificia Universidad Católica del Ecuador with an ID of QCAZ 13857. The snake was euthanized using 2% lidocaine before enucleation, following the protocol of Simmons and Muñoz-Saba (2005). After the eye was removed, the posterior eye cup was separated from the rest of ocular tissues. Dissected tissues were immersed briefly in liquid nitrogen for rapid freezing and transferred to −80°C for long-term storage.

RNA extraction

The TRIzol Plus RNA purification system (Invitrogen, Carlsbad, CA) was used for RNA extraction, using the manufacturer’s instructions. Briefly, 1 ml of TRIzol was added to frozen retinal samples and homogenized using a pestle. The samples were centrifuged at 13,000 rpm for 10 min at 4°C to separate the supernatant, which contained the RNA, from the rest of the tissue debris. The supernatant was left briefly at room temperature (RT) before adding 0.2 ml of chloroform, shaking vigorously and incubating at RT for an additional 2–3 min. Samples were centrifuged again at 13,000 rpm for 15 min at 4°C, before adding an equal volume of 70% ethanol to the aqueous phase. This mixture was then transferred to a silica column and centrifuged to bind the RNA to the membrane. The RNA was washed, treated with DNase (to remove genomic DNA and assure that any resulting amplifications are the product of solely transcript sequences) and eluted in 50 µl of nuclease-free water. The RNA yield and quality were analyzed using a Nanodrop 1000™ UV-Vis spectrophotometer (Thermo Fisher Scientific, Waltham, MA), aliquotted into 1–2 µg fractions and stored at −80°C.

Complementary DNA (cDNA) synthesis

For cDNA synthesis, the GoScript system (Promega, Madison, WI) was used, following the manufacturer’s instructions. Briefly,
1–2 μg of total RNA was reverse-transcribed using an anchored oligo-dT primer, Oligo-dT-ANC (Supplementary materials, Table 1), which also contained a specific 5’-end sequence that was utilized during the subsequent rapid amplification of cDNA ends (RACE) experiments. After each reverse transcription reaction, the reverse transcriptase enzyme was heat-inactivated and the template RNA removed by the addition of RNase H (Promega, Madison, WI). The resulting cDNA was purified using a QIAquick PCR Purification Kit (Qiagen, Germantown, MD) and eluted in 50 μl of nuclease-free water.

Polymerase chain reaction (PCR) amplification and RACE

Visual opsin coding sequences were amplified using GoTaq Flexi DNA polymerase (Promega, Madison, WI) in various nested reverse transcription PCR (RT-PCR) experiments, where both degenerate and gene-specific primers were used. The degenerate primers were designed based on known sequences of previously identified opsins (Davies et al., 2009b) to amplify the sequence of three visual opsin gene classes that are known to be expressed in snakes, namely SWS1 and LWS (both expressed in cones), and rod opsin (RH1; usually expressed in rods). Similar experiments to PCR-amplify SWS2 and RH2 (both expressed in cones), if present in B. atrox, were also conducted since these photopigments are also expressed in the retinae of reptiles. All primers were initially designed to cross multiple exon–intron boundaries so that any resulting amplification is solely the product of cDNA, thus excluding the chance of genomic DNA (gDNA) as a potential source of contamination. For the first and second rounds of PCR, the following conditions were used to amplify a core region of each opsin sequence: 94°C for 10 min (denaturation); 40 cycles at 94°C for 30 s, 45–55°C for 1 min, and 72°C for 1.5 min (amplification and extension); and a final extension at 72°C for 10 min. The nesting procedures used followed the approach outlined by Davies et al. (2009b). After the initial amplifications, RACE and gene-specific primers were used to determine the 5’- and 3’-ends of each opsin mRNA sequence: 94°C for 10 min (denaturation); 40 cycles at 94°C for 30 s, 45–55°C for 1 min, and 72°C for 1.5 min (amplification and extension); and a final extension at 72°C for 10 min. The nesting procedures used followed the approach outlined by Davies et al. (2009b). To obtain the 3’-end, oligo-dT primed cDNA was used in nested PCR amplification experiments in conjunction with gene-specific forward primers and 3’-RACE-OUT-R (i.e., OUTER) and 3’-RACE-IN-R (i.e., INNER) reverse primers (see Supplementary materials, Table 1 for details) (Davies et al., 2009a,b). By contrast, to amplify the 5’-end of each opsin mRNA sequence, terminal deoxynucleotidyl transferase (TdT) was used to add a poly-C tail to the 5’-end of the cDNA template. For each subsequent nested amplification, an anchor primer, 5’-RACE-ANC-F, which

![Fig. 1.](https://bioweb.bio/portal/QCAZ/Especimen/57289) via ID of QCAZ 13857, with permission granted courtesy of Diego Quirola (A) and Juan Carlos Sanchez (B and C).
annealed to the poly-C tail, and a gene-specific reverse primer were used (Supplementary materials, Table 1) (Davies et al., 2004a,b). RACE PCR cycling conditions were identical to those used during the RT-PCR experiments. In all cases, amplicons of the correct size were excised from 1.2% agarose gels, purified using a QIAquick Gel Extraction Kit (Qiagen, Germantown, MD) following the manufacturer’s protocol, and sequenced directly (Macrogen, Seoul, Republic of Korea).

**Molecular phylogenetic analysis**

A codon-matched nucleotide sequence alignment of 77 gnathostome (jawed vertebrate) opsin coding regions, ranging from teleosts to mammals, was generated by ClustalW (Higgins et al., 1996) and manually manipulated to refine the accuracy of cross-species comparison. Specifically, the alignment incorporated the opsin sequences of *B. atrox* (the common lancehead) (accession numbers MH244490-MH244492) compared with those expressed in the retinae of the royal python (*Python regius*) (Davies et al., 2009b), the sunbeam snake (*Xenopeltis unicolor*) (Davies et al., 2009b), and the garter snake (*Thamnophis proximus*) (Schott et al., 2016). All five opsin classes were included, with vertebrate ancient (VA) opsin sequences (Davies et al., 2010) used collectively as an outgroup given that this opsin type is a sister clade to all five visual photopigment classes. Phylogenetic analysis of 1000 replicates was conducted in MEGA6 (Tamura et al., 2013), with evolutionary history being inferred by using the maximum likelihood method based on the Tamura-Nei model (Tamura & Nei, 1993), resulting in a tree with a highest log likelihood of −39862.9953. The percentage of trees (>50%) in which the associated taxa clustered together is shown next to the branches. Initial trees for the heuristic search were obtained using the maximum likelihood method based on the Tamura-Nei model (Tamura & Nei, 1993), resulting in a tree with a highest log likelihood of −39862.9953. The percentage of trees (>50%) in which the associated taxa clustered together is shown next to the branches.

### Table 1. Experimentally determined and mean spectral peaks of absorbance for SWS1, LWS, and RH1 photopigments expressed in the retinae of the royal python (*Python regius*), the sunbeam snake (*Xenopeltis unicolor*), the garter snake (*Thamnophis proximus*), as calculated by microspectrophotometry (MSP) and UV-Vis spectrophotometric regeneration (Regen.), and those predicted for *B. atrox* (the common lancehead)

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<th>SWS1</th>
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<td>Python</td>
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<td>Garter snake</td>
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<td>Common lancehead</td>
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*Spectral analyses*

There are >40 known amino acid tuning sites that alter the spectral peak of absorbance (λ\(_{max}\)) of the five classes of opsin-based photopigments in vertebrae. As only LWS, SWS1, and RH1 opsin-based photopigment classes are relevant to snakes in this case, SWS2 and RH2 spectral tuning will not be discussed. Using conventional numbering based on the bovine rod opsin protein sequence (accession number NP001014890), a number of studies have determined that the following 27 amino acid sites are important for the spectral tuning of RH1 photopigments: 83, 90, 96, 102, 113, 118, 122, 124, 132, 154, 163, 184, 194, 195, 207, 208, 211, 214, 253, 261, 265, 269, 289, 292, 295, 299, 300, and 317 (Chan et al., 1992; Davies et al., 2012; Davies et al., 2007; Hope et al., 1997; Hunt et al., 2001; Janz & Farrrens, 2001; Sakmar et al., 1991; Yokoyama, 2000; Yokoyama, 2008; Yokoyama et al., 1999; Yokoyama et al., 2008; Yokoyama et al., 2007). Similar studies, when applied to other photopigment classes, have determined that site 86 is critical for generating UV-sensitive (UVS) SWS1 pigments (Cowling et al., 2002), whereas five sites (namely 164, 181, 261, 269, and 292) are important for determining the λ\(_{max}\) values of LWS photopigments (Davies et al., 2012; Yokoyama, 2000). Inspection of these sites was used to estimate the λ\(_{max}\) value for the three photopigments, LWS, SWS1, and RH1, expressed in the eye of the common lancehead. The overall spectral effects of these sites and their relevance to snake vision are presented in the Results and Discussion sections below. Dark spectra for visual photopigments identified in *B. atrox* (common lancehead), *P. regius* (royal python), *X. unicolor* (sunbeam snake), and *T. proximus* (garter snake) (Davies et al., 2009b; Schott et al., 2016) were generated using a standard A\(_T\)-based rhodopsin template (Govardovskii et al., 2000).

### Results

Using cDNA generated from freshly extracted *B. atrox* retinal RNA, the coding sequences for three visual opsin genes were successfully amplified, namely SWS1, LWS, and RH1. Despite numerous attempts, using both degenerate and reptile-specific primers (Davies et al., 2009b), RH2 and SWS2 sequences were not amplified, thus suggesting that these two opsins are not found in this snake species. Through a combination of RT-PCR, and 5’- and 3’-RACE, full-length coding sequences were obtained for RH1 (1059 bp) and LWS (1098 bp) opsin genes, whereas amplification of the SWS1 opsin gene yielded a partial sequence of 826
nucleotides. On closer inspection of a codon-match alignment, each sequence was observed to contain no indels (Fig. 2). Furthermore, the genes and their resultant photopigments were deemed to be functional based on intact sequence fidelity; expression in the retina of *B. atrox*; high nucleotide and amino acid sequence identity to other snake opsin sequences; and the presence of key structural and functional features, including the seven transmembrane domains (TMDs) shown for bovine rod opsin were determined by crystallography (Palczewski et al., 2000), with the seven putative TMDs for each snake photopigment being determined online using TMHMM Server Version 2.0 (http://www.cbs.dtu.dk/services/TMHMM/). In all cases, TMDs are indicated by gray shading. Residues known to be critical for both the correct opsin protein tertiary structure and biochemical function are boxed: comparison with other photopigment sequences confirmed that these sites are conserved in all other visual pigments that derive from *B. atrox*. Using the conventional numbering system of the bovine rod opsin sequence, these key sites include (1) two conserved cysteine (C) residues at positions 110 (TMD3) and 187 (ECD2) that are involved in disulphide bond formation (Karnik et al., 1988); (2) a conserved glutamate (E) at position 134 (TMD3) that provides the negative counterion to the proton of the Schiff base (Sakmar et al., 1989); (3) a conserved glutamate (E) at position 134 (TMD3) that provides a negative charge to stabilize the inactive opsin molecule (Cohen et al., 1992); (4) a conserved lysine (K) at position 296 (TM7) that is covalently linked to the chromophore in a Schiff base (Dratz & Hargrave, 1983); (5) conservation of two cysteine (C) residues at putative palmitoylation positions 322 and 323 (Ovchinnikov et al., 1988) in the RH1 photopigment of *B. atrox*, but not LWS (please note that putative palmitoylation sites for SWS1 could not be determined); (6) the presence of a number of Ser (S) and Thr (T) residues in the carboxy terminus, which are potential targets for phosphorylation by rhodopsin kinases in the deactivation of metarhodopsinII (Palczewski et al., 1993; Zhao et al., 1997) (putative phosphorylation sites for LWS photopigments could not be determined); and (7) the conserved glycosylation sites at positions 2 and 15 (Kaučuk et al., 1994) in the RH1 opsin identified in the retina of the lancehead. Amino acids important for the spectral tuning of LWS, SWS1, and RH1 visual pigments are indicated in Figs. S1–S3.

Phylogenetic analysis was performed using a maximum likelihood approach on a codon-matched codon sequence alignment, which comprised the three common lancehead opsin sequences (accession numbers MH244490-MH244492) compared with the coding regions of 77 gnathostome sequences from all five opsin classes, namely LWS, SWS1, SWS2, RH2, and LWS. The neighbor-joining phylogenetic tree shown in Fig. 3 clearly indicated that each *B. atrox* opsin sequence clades within the orthologous pigment gene classes of *LWS, SWS1,* and *RH1,* which were accompanied by high bootstrap values at each node. In particular, the opsins expressed in the retina of *B. atrox* were robustly clustered together with other snake sequence used in the analysis, namely the royal python (*Python regius*) (Davies et al., 2009b), the sunbeam snake (*Xenopeltis unicolor*) (Davies et al., 2009b), and the garter snake (*Thamnophis proximus*) (Schott et al., 2016). In all cases, the common lancehead opsins were more closely related to the garter snake than the two “lower order” henophidians, thus reflecting the shared evolutionary history between higher order” snake species within Alethinophidia.

To predict RH1 spectral tuning, 27 known tuning sites were compared between the predicted *B. atrox* and bovine rod opsin protein sequences, namely amino acids 83, 90, 96, 102, 113, 118, 122, 124, 132, 164, 183, 194, 195, 207, 208, 211, 214, 253, 261, 265, 269, 289, 292, 295, 299, 300, and 317 (Chan et al., 1992; Davies et al., 2012; Davies et al., 2007; Hope et al., 1997; Hunt et al., 2001; Janz & Farrens, 2001; Sakmar et al., 1991, 2000, 2008; Yokoyama et al., 2007; Yokoyama et al., 2008; Yokoyama et al., 2009). With a complement of Asp-Gly-Tyr-Tyr-Glu-Thr-Glu-Ala-Asp-Thr-Ala-Lys, which are potential targets for phosphorylation by rhodopsin kinases in the deactivation of metarhodopsinII (Palczewski et al., 1993; Zhao et al., 1997) (putative phosphorylation sites for LWS photopigments could not be determined); and (7) the conserved glycosylation sites at positions 2 and 15 (Kaučuk et al., 1994) in the RH1 opsin identified in the retina of the lancehead. Amino acids important for the spectral tuning of LWS, SWS1, and RH1 visual pigments are indicated in Figs. S1–S3.

![Fig. 2. A codon-matched alignment of the amino acid sequences of RH1, LWS, and SWS1 expressed in the retina of *B. atrox*. Bos taurus (BT) rod opsin (RHK) is used as a reference. Each residue is color-coded based on their biochemical properties (e.g., charged vs. hydrophobic), with asterisks denoting identical consensus residues between all snake opsin sequences and bovine RH1. Dashes represent gaps that were inserted to maintain a high degree of sequence identity present between the different opsins classes. The transmembrane domains (TMDs) shown for bovine rod opsin were determined by crystallography (Palczewski et al., 2000), with the seven putative TMDs for each snake photopigment being determined online using TMHMM Server Version 2.0 (http://www.cbs.dtu.dk/services/TMHMM/). In all cases, TMDs are indicated by gray shading. Residues known to be critical for both the correct opsin protein tertiary structure and biochemical function are boxed: comparison with other photopigment sequences confirm that these sites are conserved in all three visual pigments that derive from *B. atrox*. Using the conventional numbering system of the bovine rod opsin sequence, these key sites include (1) two conserved cysteine (C) residues at positions 110 (TMD3) and 187 (ECD2) that are involved in disulphide bond formation (Karnik et al., 1988); (2) a conserved glutamate (E) at position 134 (TMD3) that provides the negative counterion to the proton of the Schiff base (Sakmar et al., 1989); (3) a conserved glutamate (E) at position 134 (TMD3) that provides a negative charge to stabilize the inactive opsin molecule (Cohen et al., 1992); (4) a conserved lysine (K) at position 296 (TM7) that is covalently linked to the chromophore via a Schiff base (Dratz & Hargrave, 1983); (5) conservation of two cysteine (C) residues at putative palmitoylation positions 322 and 323 (Ovchinnikov et al., 1988) in the RH1 photopigment of *B. atrox*, but not LWS (please note that putative palmitoylation sites for SWS1 could not be determined); (6) the presence of a number of Ser (S) and Thr (T) residues in the carboxy terminus, which are potential targets for phosphorylation by rhodopsin kinases in the deactivation of metarhodopsinII (Palczewski et al., 1993; Zhao et al., 1997) (putative phosphorylation sites for LWS photopigments could not be determined); and (7) the conserved glycosylation sites at positions 2 and 15 (Kaučuk et al., 1994) in the RH1 opsin identified in the retina of the lancehead. Amino acids important for the spectral tuning of LWS, SWS1, and RH1 visual pigments are indicated in Figs. S1–S3.](https://www.cambridge.org/core/terms)
Fig. 3. Phylogenetic analysis of nucleotide sequences derived from B. atrox (the common lancehead; accession numbers MH244490-MH244492), LWS, SWS1, and RH1 opsin genes, compared with those determined for the royal python (Python regius) (Davies et al., 2009), the sunbeam snake (Xenopeltis unicolor) (Davies et al., 2009), and the garter snake (Thamnophis proximus) (Schott et al., 2016). These snake sequences were compared to other vertebrate visual pigment coding sequences for all five visual opsin classes, namely LWS, SWS1, SWS2, RH2, and RH1. All sequences were used to generate a codon-matched alignment with vertebrate ancient (VA) opsin sequences (Davies et al., 2010) used collectively as an outgroup given that this opsin type is a sister clade to all five visual photopigment classes. Posterior probability values (represented as a percentage, where only those greater than 50% are shown) are indicated for each resolved node. The scale bar indicates the number of nucleotide substitutions per site within each branch length. The sequences used for generating the tree are as follows: (1) RH1 opsin class: human (Homo sapiens), NM000539; mouse (Mus musculus), NM145383; fat-tailed dunnart (Sminthopsis crassicaudata), AY159786; platypus (Ornithorhynchus anatinus), EF050076; chicken (Gallus gallus), NM205490; pigeon (Columba livia), AH007730; green anole (Anolis carolinensis), AOITRHOOPS; sunbeam snake (Xenopeltis unicolor), FJ497233; python (Python regius), FJ497236; garter snake (Thamnophis proximus), KU306726; common lancehead (Bothrops atrox), MH244492; African clawed frog (Xenopus laevis), NM001087048; Comoran coelacanth (Latimeria chalumnae), AF131253; Australian lungfish (Neoceratodus forsteri), EF526295; goldfish (Carassius auratus), L11863; zebrafish (Danio rerio), NM131084; (RH1.1); (2) RH2 opsin class: chicken (Gallus gallus), NM205490; pigeon (Columba livia), AH007731; green anole (Anolis carolinensis), AH004781; Comoran coelacanth (Latimeria chalumnae), AF131258; Australian lungfish (Neoceratodus forsteri), EF526296; goldfish (Carassius auratus), L11863; zebrafish (Danio rerio), NM131253; (RH2.1); (2) SWS2 class: platypus (Ornithorhynchus anatinus), EF050077; chicken (Gallus gallus), NM205517; pigeon (Columba livia), AH007799; green anole (Anolis carolinensis), AF133907; African clawed frog (Xenopus laevis), BC080123; Australian lungfish (Neoceratodus forsteri), EF526295; goldfish (Carassius auratus), L11863; zebrafish (Danio rerio), NM131253.
et al., 2008). For B. atrox, Ala is found at site 292 (underlined), so Thr195 will not affect spectral tuning; as such, the predicted $\lambda_{\text{max}}$ value for the RH1 photopigment in the common lancehead is 500 nm (i.e., identical to the bovine rod photopigment (Hubbard, 1969) (Fig. 4).

For SWS1 and LWS photopigments, predictions of the spectral peaks of absorbance are far less complicated than that of RH1 photopigments. For example, the presence of PhE86 in the predicted sequence for the SWS1 photopigment of B. atrox means that it will be UVS, with a $\lambda_{\text{max}}$ value at 360 nm (Cowing et al., 2002) (Fig. 4). For LWS photopigments, a complement of Ala\textsuperscript{164}-His\textsuperscript{181}-Tyr261-Thr269-Ala292 in B. atrox only differs from the typical vertebrate complement of Ser\textsuperscript{164}-His\textsuperscript{181}-Tyr261-Thr269-Ala292 at one site. As such, the presence of a Ser\textsuperscript{164}Ala substitution (highlighted in bold) in the predicted LWS sequence of the common lancehead will shift the $\lambda_{\text{max}}$ value from 560 nm (Davies et al., 2012; Yokoyama, 2000) by 7 nm (Davies et al., 2012; Yokoyama, 2000; Yokoyama, Yang, & Starmer, 2008) to 553 nm (Fig. 4). In summary, the $\lambda_{\text{max}}$ values for the three functional photopigments expressed in the retina of B. atrox are predicted to be 360 nm, 500 nm, and 553 nm for SWS1, RH1, and LWS opsin proteins, respectively (Fig. 4).

Discussion

In this study, three photopigment genes were identified as being expressed in the retina of Bothrops atrox, namely SWS1 (cone), LWS (cone), and RH1 (rod). By contrast, the other two opsin classes, namely SWS2 and RH2, which are expressed in many other reptiles, but have never been identified in snakes, were not identified. This is consistent with the likely loss of these genes in the ancestor to all snakes, including their omission from two sequenced snake genomes (Castoe et al., 2013; Simões et al., 2015; Simões & Gower, 2017; Vonk et al., 2013). This is an identical complement of opsin genes to both lower order (i.e., more ancestral) and “higher order” snakes, except for the burrowing blind snakes of the Order Scodichelida, where only RH1 is present (Davies et al., 2009b; Simões et al., 2015). The absence of expression of these opsins is believed to be the result of the nocturnal and fossorial nature of many snakes, and presumably of the ancestral snake as well (Davies et al., 2009b). However, as the complement of pigment genes is the same as many mammals (excluding monotremes and some trichromatic primates), it is possible that ancestral snakes were subjected to a “mesopic bottleneck” instead of a restricted nocturnal phase, where it may be predicted that all cones would be lost; this hypothesis was first proposed to explain the pigment complement in the mammalian lineage (Davies et al., 2012), but is also relevant to snake evolution (Davies et al., 2012; Davies et al., 2009b). In this investigation, visual opsin genes were identified from the first ever South American viper, where the resultant predicted protein sequences were used to calculate the spectral peaks of absorbance (i.e., the $\lambda_{\text{max}}$ values) for B. atrox.

B. atrox is a nocturnal predatory snake found in tropical forest grounds, where even moonlight and/or starlight is restricted by the dense forest foliage. The exact retinal morphology and rod-to-cone ratio are unknown in the B. atrox retina; as with many nocturnal animals, and given the common lancehead’s predatory behavior, the visual system of B. atrox may be under strong selective pressures to maximize photon absorption in dim-light environments (Cummins & Partridge, 2001; Lythgoe, 1984; Partridge &...
Cummings, 1999). Individuals belonging to this species need to move, hunt, and mate in nocturnal conditions (with restricted moonlight and/or starlight), which could explain their need for a more rod-rich retina and the presence of only a small population of cones.

Even under different nocturnal light environments, large differences may exist in terms of opened or closed canopies, foliage density, lunar phases, cloud cover, etc. These differences could greatly modify the amount of light available (Lythgoe, 1972; Munz & McFarland, 1973; Pariente, 1980). Vipers such as the African viper *Echis ocellatus* and South American common lancehead *B. atrox* inhabit rainforest grounds, where light can be extremely limited because of the dense foliage. This is likely to change the selective pressures acting upon the visual system of these snakes compared with other animals, especially those that inhabit more open environments with less foliage, for example grasslands and savannas where the African vipers *Causus rhombeatus* and *Bitis nasicornis* are found. Rainforests are also richer in longer wavelengths (i.e., red-shifted) than dry forest and grasslands (Veilleux & Cummings, 2012), which could be another reason why *B. atrox* and *E. ocellatus* visual photopigments are predicted to be more long wavelength-shifted in comparison with those of other snakes (Simões et al., 2016a).

In this study, the spectral peaks for *B. atrox* photopigments were calculated to be 360 nm, 500 nm, and 553 nm for SWS1, RH1, and LWS, respectively. Unlike SWS2 and RH2, which require extensive *in vitro* regeneration of pigments and the spectral shifts associated with mutant opsin constructs *via* site-directed mutagenesis (Bowmaker, 2008; Davies et al., 2012; Yokoyama, 2000), these three photopigments are so well studied that it is possible to predict their $\lambda_{max}$ values solely from their protein sequence, without the need for further *in vitro* regeneration of MSP experiments.

Comparison with henophidian snakes (e.g., the sunbeam snake and the python) shows similarity in their spectral absorbance maxima [i.e., on average 361 nm (SWS1), 496 nm (RH1), and 554 nm (LWS)]. By contrast, the garter snake possesses cone photopigments with $\lambda_{max}$ values that are shifted at either end of the light spectrum [i.e., 365 nm for SWS1 (i.e., $+5$ nm) and 544 nm for LWS (i.e., $-10$ nm)], thus decreasing the range of wavelengths being detected. As such, the photopigments of *B. atrox* do not resemble other phylogenetically relevant “higher order” snakes, but are similar to “lower order” species from a spectral perspective. This suggests that the visual system of *B. atrox* has adapted to a particular habitat and not a phylogenetic remnant, since *B. atrox* and “lower order” snakes generally live in dim-light or scotopic environments resembling other nocturnal vertebrates (Simões et al., 2016a; Veilleux & Cummings, 2012), whereas the garter snake is diurnal and possesses a pure-cone retina, despite the expression of the rod (RH1) opsin classes.

Compared with RH1 and LWS, where expression levels were relatively high enough to obtain full-length sequences, *B. atrox* SWS1 was more problematic. Indeed, it was not possible to obtain a full-length sequence, which could be due to the relatively low expression within the retinæ of nocturnal vertebrates (Ahnelt & Kolb, 2000; Davies et al., 2009b; Hunt & Peichl, 2014), or, alternatively, the fact that although many primers were designed to amplify this gene, none of them was adequate to amplify its full length. As RACE experiments were able to extend the sequence a little further to result in a protein sequence that did not contain indels or spurious mutations when compared to other vertebrate SWS1 photopigments, it is concluded that SWS1 is likely to be functional in this snake species. This was corroborated by the fact that the long partial SWS1 protein sequence contained all the expected residues required for structural conformity and spectral tuning. It should be noted here that in spite of the fact that the SWS1 gene was not cloned in its entirety, the spectral peak of absorbance of the respective protein could be predicted as the sequence did include amino acid 86, which has been shown to determine SWS1 spectral tuning almost exclusively (Cowling et al., 2002; Fasick et al., 2002). In general, animals possess far less UV or blue cones than rods, red cones, or green cones and, therefore, the corresponding SWS1 gene and protein have been found to be expressed at significantly lower levels. It would be of interest to determine whether SWS1 sensitivity to UV wavelengths is correlated with a nocturnal lifestyle within snakes. However, since the diurnal garter snake also possesses a UVS cone (Schott et al., 2016), the correlation between UVS photopigments and nocturnality may not be a general observation.

To obtain a more complete appreciation of how visual adaptation is associated with both the habitat and activity patterns of animals, *B. atrox* was compared to further examples of nocturnal and diurnal snakes (Simões et al., 2016a). *Oxyrhopus melanogenys* is a nocturnal colubrid that occupies a similar habitat as *B. atrox* and possesses photopigments with $\lambda_{max}$ values similar to those of *B. atrox* (Simões et al., 2016a). By contrast, *Spalerosophis diadema* is a diurnal colubrid that inhabits more open locations without a canopy (such as grasslands and deserts). In *S. diadema*, the $\lambda_{max}$ value of the RH1 photopigment is ~20 nm short-wavelength-shifted, whereas the SWS1 cone is likely to be violet sensitive (VS) and not UVS like in nocturnal snakes. This suggests that a narrower spectral range may be common in diurnal snake vision [e.g., ~400 nm to 555 nm for *S. diadema* (Simões et al., 2016a); 365 nm to 544 nm for the garter snake (Schott et al., 2016)] compared with the spectral range of vision in nocturnal species (360 nm to 554 nm) (Davies et al., 2009b). These spectral differences conform to the view that the habitat of an animal, as well as its behavior, plays a crucial role in the evolution of the visual system.

For henophidian snakes that are at least partly fossorial, crepuscular, or nocturnal, it was proposed that the python and the sunbeam snake could be dichromatic based on the identification of two cone photopigments (Davies et al., 2009b). However, the spectral overlap between UVS SWS1 cones and LWS cones is minimal so it was thought that they would not be involved in color vision. Physiological evidence has shown that under mesopic conditions, where both rods and cones would be active (Buck, 1997; Reitner et al., 1991; Stabell & Stabell, 1994), the presence of three photopigments [SWS1, LWS, and the rod (RH1) opsins] that spectrally overlapped could confer a rudimentary color visual system (a form of conditional trichromacy) (Davies et al., 2009b) to these species, despite the large spectral range covered by the outermost cone types (Davies et al., 2009b; Sillman et al., 1999; Sillman et al., 2001). As a nocturnal species, *B. atrox* is not expected to possess color vision as at night (and under dense foliage that diminishes moonlight and starlight) both SWS1- and LWS-expressing cones should be inactive given their threshold for activity in bright-light conditions. Nonetheless, it has been known for the common lancehead to be occasionally partly active at twilight and during the day, especially when hunting is a necessity (Oliveira & Martins, 2001). As rods and cones are active at dusk and dawn, the presence of three functional photopigments in *B. atrox* may also confer conditional trichromacy and a selective advantage when hunting prey.
Future work

In this work, the visual system was investigated for B. atrox, the common lancehead, one of the most poisonous snakes in the Amazon region (Oliveira & Martins, 2001). More specifically, the visual opsins expressed in this viper were studied. However, many questions still remain unanswered, especially considering the transmutation theory proposed by Walls (Walls, 1934, 1942). It would be of great interest, for example, to examine the potential transmutation of B. atrox photoreceptors at the genetic level, something that has never been examined in a viper. To assess this, a future study could be conducted that investigates the different isoforms of the visual phototransduction cascade genes that are present and functional in the retina of the common lancehead, such as GNAT, retGC, arrestins, transducins, among others. Additionally, it would be of interest to assess photoreceptor morphology in detail using electron microscopy to visualize the different intermediate morphotypes present, if any.

Also, it would be of interest to investigate the developing visual system of B. atrox, to determine whether it is different at diverse developmental time points (e.g., embryo vs. juvenile vs. adult). Such differences have never been demonstrated in reptiles; however, it is well known that in some mammals, such as rodents, all cones start as S-cones or short-wavelength-sensitive cones and then differentiate to become L-cones or long-wavelength-sensitive cones (Lucats et al., 2005; Széll et al., 1994). It would be fascinating to discover if reptiles follow the same pattern or are more similar to other mammals like humans who express the same opsins differences have never been demonstrated in reptiles; however, it is of interest to study whether there are sex-dependent differences in opsin expression in B. atrox, it would be of interest to study the visual ecology of basal vertebrates. Molecular Biology and Evolution 26, 1803–1809.

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